

# *Heleobia dobrogica* (Grossu & Negrea, 1989)(Gastropoda: Rissooidea: Cochliopidae) and the estimated time of its isolation in a continental analogue of hydrothermal vents

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### Abstract

*Heleobia dobrogica* is the only gastropod species living in the Movile Cave in Dobrogea, Romania. In the cave there is little oxygen but large amounts of carbon dioxide and methane in the atmosphere, and a large amount of hydrogen sulphide in the water. All the non-predatory animals feed on the chemoautotrophic microorganisms that draw energy from the sulphide hot springs beneath the cave. Five COI mtDNA sequences were used for maximum likelihood phylogeny reconstruction together with eight sequences of cochliopids and two outgroup rissooids from GenBank., *Salenthydrobia ferrerii* and *Peringia ulvae* were used as an outgroup for the calibration of the molecular clock. The estimated time of divergence between the two species was 2.172±0.171 Mya. This coincides with the period marking the beginning of the fall in temperature and precipitation that initiated the glacial period in Europe, predating the Pleistocene. Most probably at that time *Heleobia dobrogica* found a safe shelter within a warm cave. Our results suggest that *H. dobrogica* is closely related to *H. dalmatica*, and both species may be congeneric with *Heleobops docimus*.

Key words: troglobiont, Cochliopidae, phylogeny, mtDNA, molecular clock, Romania

## Introduction

In 1986, while excavating for a construction project, engineers found an unusual cave near Mangalia (south of Constanta in Dobrogea), a few kilometres from the Black Sea coast. The small cave (12,000 m<sup>2</sup>), named Movile, is about 300 metres long and less than 3 metres high. It contains a small lake in its lower part. The physical and chemical conditions within the cave are unusual: the water is rich in hydrogen sulphide (8–12 mg/l); and the atmosphere is poor in oxygen (7–10%), rich in carbon dioxide (2–3.5%) and charged with a significant amount of methane (1-2%)(Marin and Nicolescu 1993; Sarbu et al. 1996). The cave has no natural entrance and the only man-made entrance is entirely sealed off (by two gates and an airtight lid) to prevent alteration of the natural conditions within the cave. Forty-six invertebrate species, including 33 endemics, have been found in the cave: 28 of these (22 endemic) are terrestrial and 18 (11 endemic) are aquatic, one gastropod included. All of the primary consumers in this ecosystem feed on the chemoautotrophic bacteria and fungi that draw energy from the sulphide hot springs beneath the cave (Sarbu 1991; Sarbu and Kane 1995; Sarbu et al. 1996; Lascu 2001). This is a clear analogue to the well-known hydrothermal vents ('hot vents') found in oceanic rift zones. Lascu (1989) hypothesized that this unusual fauna found a shelter in the cave some five million years ago, when the climate became colder.

In 1989 Grossu and Negrea described a new species of *Paladilhiopsis* Pavlovic, 1913 from the lake in Movile Cave. Bernasconi (1991) subsequently transferred this species to the genus *Heleobia* Stimpson, 1865 (*=Semisalsa* Radoman, 1974) and studied the shell variation and anatomy of this species (Bernasconi 1994, 1997), while Szarowska (2006) described and illustrated its protoconch and female reproductive anatomy. The shells of *Heleobia dobrogica* are illustrated in Fig. 1.



**FIGURE 1:** Shells of *Heleobia dobrogica*: A,B—female, C,D—male. Movile Cave, ZMUJ RO06M1, 2, 3, 4, respectively.

The aim of the present paper is to try to estimate the time of isolation of *Heleobia dobrogica* (Grossu & Negrea, 1989) in the cave, as a probable time of isolation of all the Movile Cave fauna, applying the molecular clock approach. We also intend to infer the relationships of *H. dobrogica* within the Cochliopidae.

## Material and methods

Snails were collected by hand, fixed with 80% ethanol and stored in 96% ethanol. They were hydrated in TE buffer (3 x x

10 min.), DNA was extracted with the SHERLOCK extracting kit (A&A Biotechnology), and the final product was dissolved in 20 µm of TE buffer. The PCR reaction was performed with the following primers: LCOI490 (5'-GGTCAACAAATCATAAAGATATTGG-3') and COR722b (5'-TAAACTTCAGGGTGACCAAAAAATYA-3') for the COI gene (Folmer et al. 1994). The PCR conditions were as follows: 4 min at 94 °C followed by 35 cycles of 1 min at 94 °C, 1 min. at 55 °C, 2 min. at 72 °C, with an additional elongation step of 4 min performed at 72 °C after all the cycles. The total volume of each PCR reaction mixture was 50 µl. 10 µl of the PCR product was run on 1 % agarose gel to check the quality of the PCR products. The PCR product was purified using Clean-Up columns (A & A Biotechnology). The purified PCR product was sequenced (Hillis et al. 1996a) using BigDye Terminator v3.1 (Applied Biosystems) and the protocols and primers described above. The sequencing reaction products were purified using ExTerminator Columns (A & A Biotechnology), and the sequences were read using the ABI Prism sequencer.

The sequences were aligned by eye, using BioEdit 5.0.0 (Hall 1999) and edited with MacClade 4.05 (Maddison and Maddison 2002).

The only sequence of Heleobia available was the H. dalmatica (Radoman, 1974) found in GenBank, thus we used it for the estimation of time of isolation. All the cochlipid sequences available from GenBank were used for the phylogenetic analysis. The sequences of Radomaniola callosa (Paulucci, 1881) (Hydrobiidae) and Bithynia tentaculata (Linnaeus, 1758) (Bithyniidae) were included, as a multiple outgroup. To test the molecular clock, the same set of cochliopid taxa was used, the outgroups were the hydrobiids Peringia ulvae (Pennant, 1777) and Salenthydrobia ferrerii Wilke, 2003 whose divergence time was used to calibrate the clock.

**TABLE 1.** GenBank Accession Numbers and references of the COI sequences of species considered (*Heleobia dobrogica*: specimens ZMUJ RO06M10- ZMUJ RO06M14, respectively).

Species	GenBankAN	references	extraction
Heleobia dobrogica (Grossu & Negrea, 1989) <sup>1</sup>	EU938128	present study	M5R
Heleobia dobrogica (Grossu & Negrea, 1989) <sup>2</sup>	EU938129	present study	E09
Heleobia dobrogica (Grossu & Negrea, 1989) <sup>3</sup>	EU938130	present study	AB11
Heleobia dobrogica (Grossu & Negrea, 1989) <sup>4</sup>	EU938131	present study	G115
Heleobia dobrogica (Grossu & Negrea, 1989) <sup>5</sup>	EU938132	present study	G116
Heleobia dalmatica (Radoman, 1974) 1	AF367631	Wilke et al. 2001	
Heleobia dalmatica (Radoman, 1974) 2	AF129321	Hershler et al. 1999	
Heleobops docimus Thompson, 1968	AF129322	Hershler et al. 1999	
Onobops jacksoni (Bartsch, 1953)	AF367645	Wilke et al. 2001	
Spurwinkia salsa (Pilsbry, 1905),	AF367633	Wilke et al. 2001	
Cochliopa sp.	AF354762	Liu et al. 2001	
Pyrgophorus platyrachis Thompson, 1968	AF367632	Wilke et al. 2001	
Littoridinops monroensis (Frauenfeld, 1863)	AF367644	Wilke et al. 2001	
Radomaniola callosa (Paulucci, 1881)	AF367649	Wilke et al. 2001	
Bithynia tentaculata (Linnaeus, 1758)	AF367643	Wilke et al. 2001	
Salenthydrobia ferrerii Wilke, 2003	AF449200	Wilke 2003	
Peringia ulvae (Pennant, 1777)	AF118288	Wilke 2003	

The maximum likelihood (ML) approach often tends to find the wrong reconstructions, especially in analyses involving a large number of taxa with short sequences (Nei *et al.* 1998; Nei and Kumar 2000). There is no parameter associated with a tree topology in the entire maximum likelihood theory: one must simply assume that the tree with the "truest" branch lengths is also the one with the best topology (Yang *et al.* 1995; Nei 1987, 1996). There is also strong evidence that the more complicated the model of evolution, the higher the variance of the resulting reconstructions (Nei and Kumar 2000). Our knowledge of the evolution of DNA is incomplete, thus all the available models are probably unrealistic. Thus, it may happen that the simplest models will result in phylogeny reconstructions which are closest to the real historical processes (Gaut and Lewis 1995; Yang 1997; Takahashi and Nei 2000, Falniowski 2003). On the other hand, similar remarks may be made about other phylogenetic techniques as well, and the ML approach is not sensitive to the violation of some of its assumptions (Swofford *et al.* 1996). Thus we decided to apply the maximum likelihood approach as implemented in PAUP\*4.0b10 (Swofford 2002). PAUP together with Modeltest (Posada and Crandall 1998) was used to find the appropriate model of evolution, with the Akaike Information

Criterion (Posada and Buckley 2004). This model was also selected for the set of taxa with *Peringia* and *Salenhydrobia* as an outgroup, and the best ML trees were found to perform the Likelihood Ratio Test (LRT) (Nei and Kumar 2000; Posada 2003) with PAUP. MEGA4 (Tamura *et al.* 2007) was used to run the Relative Rate Tests (RRT) (Tajima 1993). The pairwise ML distances were calculated with PAUP. Wilke's (2003) data were used to calibrate the clock. Maximum Composite Likelihood ( $\Gamma$ ) with standard errors (1000 bootstrap replicates) was calculated with MEGA4.

The specimens are lodged in the collection of the Department of Malacology of Jagiellonian University, Kraków (ZMUJ RO06M).

#### Results

Molecular distances and estimation of time of divergence

Five partial (638 bp) sequences of COI (Table 1) represented four haplotypes that differed in seven positions. For all the cochliopid taxa, with Salenthydrobia ferrerii and Peringia ulvae as outgroups, the Akaike Information Criterion (AIC) selected the model TIM+I+G, with base frequencies: A = 0.2678, C = 0.1459, G = 0.1627, T = 0.4236, substitution rate matrix: [A-C] = 1.0000, [A-G] = 537.9000, [A-T] = 59.6603, [C-G] = 59.6603, [C-T] =285.8066, [G-T] = 1.0000, proportion of invariable sites: (I) = 0.6183, and  $\Gamma$  distribution with the shape parameter 1.5631. The LRT of this data set does not reject the molecular clock hypothesis (log  $L_0 = -2733.4536$ , log  $L_1 =$ -2719.8802, Δ = 27.1468, DF = 13, P>0.1239. Tajima's RRT for Heleobia *dobrogica* and *H. dalmatica* with Salenthydrobia and Peringia as an outgroup resulted in P>0.2230, and P>0.7237, respectively, thus not rejecting the molecular clock. RRTs for each pair of cochliopids with either Salenthydrobia or Peringia as an outgroup did not reject the molecular clock hypothesis either, except where Onobops was included and Salenthydrobia was used as an outgroup. Thus Onobops jacksoni (Bartsch, 1953) was excluded from the data set, for which the model GTR+I+G was selected, with base frequencies: A = 0.2682, C = 0.1526, G = 0.1627, T = 0.4166, substitution rate matrix: [A-C] = 0.0005, [A-G] = 20.4928, [A-T] = 1.6171, [C-G] = 2.1206, [C-T] = 9.0843, [G-T] = 1.0000, proportion of invariable sites: (I) = 0.6343, and  $\Gamma$  distribution with the shape parameter 2.5596. The LRT of this data set also does not reject the molecular clock hypothesis (log  $L_0 = -2546.0733$ ,  $\log L_1 = -2538.6285$ ,  $\Delta = 14.8874$ , DF = 12, P>0.2940.

The pairwise ML distances calculated for both models are given in Table 2. The distances calculated for the model without *Heleobops* (0.05774-0.07135, mean 0.0636 $\pm$ 0.005) were used to estimate the time of divergence. The value 0.15605 between *Salenthydrobia* and *Peringia* was calibrated for 5.33 Mya by Wilke (2003). Thus, applying this calibration, the mean divergence time between the two species of *Heleobia* was 2.172 $\pm$ 0.171 Mya (2.139 $\pm$ 0.167 Mya for the model calculated including *Onobops jacksoni*). The Maximum Composite Likelihood distances (Г TABLE 2. Pairwise distances: below diagonal: for the model selected for the cochliopids with Salenthydrobia ferrerii and Peringia ulvae as outgroup; above diagonal: for the model selected

distribution,	α=1.5631)	gave	estimates	within	the	range
$2.390{\pm}0.665$	Mya – 2.870	0±0.73	33Mya.			

	1 2	m	4	ம	9	7	ø	ი	10	11	12	13	14	15
1. Heleobia dobrogica 1	0.0046	10.00943 0	.006210.	006210.	07135 0.	07135 0.	08285 -	.0	45423 0.	570450.	570010.	.45233 1	.062880	93075
2. Heleobia dobrogica 2	0.00463	- 0.00463 0	.00463 0.	00463 0.	.064310.	06431 0.	08023 -	. 0	451570.	552560.	54563 0.	.433791	.03172 0.	91376
3. Heleobia dobrogica 3	0.00941 0.0046	2 0	.00621 0.	006210.	05774 0.	05774 0.	07794 -	. 0	426990.	530510.	51499 0.	.400151	.064210.	94848
4. Heleobia dobrogica 4	0.00619 0.0046	2 0.00621 -	. 0	000000	.062300.	06230 0.	07740 -	. 0	428170.	529700.	51073 0.	.405371	.052550.	92934
5. Heleobia dobrogica 5	0.00619 0.0046	2 0.00621 0	00000.	0	.062300.	06230 0.	07740 -	.0	428170.	529700.	51073 0.	.405371	.052550.	92934
6. Heleobia dalmatica l	0.07183 0.0647	9 0.05821 0	.062760.	06276 -	.0	000000.	07135 -	.0	402600.	523970.	537070.	.403801	.057760.	98889
7. Heleobia dalmatica 2	0.07183 0.0647	9 0.05821 0	.062760.	06276 0.	00000.	0.	07135 -	.0	406700.	523970.	45594 0.	.428891	.057760.	94705
8. Heleobops docimus	0.08523 0.08259	9 0.08027 0	.079870.	079870.	07541 0.	07541		. 0	402600.	54283 0.	537070.	.403801	.063010.	98889
9. Onobops jacksoni	0.29101 0.2804	2 0.26942 0	.26972 0.	26972 0.	34971 0.	34971 0.	28906 -	• • • • • • • • •						
10. Spurwinkia salsa	0.40012 0.3984	3 0.38311 0	.38033 0.	38033 0.	.39270 0.	39270 0.	38580 0	.26464	0.	283290.	13245 0.	.065891	.077291.	32330
11. Cochliopa sp.	0.60086 0.58128	3 0.56351 0	.55892 0.	55892 0.	57809 0.	57809 0.	60960 0	.38109 0.	31285	0.	32152 0.	.262911	.07483 1.	13441
12. Pyrgophorus platyrachis	0.53188 0.5092	2 0.48418 0	.47586 0.	47586 0.	54765 0.	54765 0.	44527 0	.353410.	139300.	35811		.123771	.327971.	30126
13. Littoridinops monroensis	0.44204 0.4243	7 0.39528 0	.39667 0.	39667 0.	42292 0.	42292 0.	44038 0	.30130 0.	070070.	288400.	13025	1	.11022 1.	30494
14. Salenthydrobia ferrerii	0.91425 0.8957	5 0.93218 0	.91853 0.	91853 0.	96271 0.	96622 0.	96271 0	.893611.	288631.	205401.	398631.	.36799 -	0	15605
15. Peringia ulvae	1.03326 0.9963	0 1.03949 1	.02609 1.	02609 1.	06461 1.	07114 1.	06461 0	.884461.	033351.	149621.	338961.	.15377 0	.15968 -	

Phylogenetic analysis

The Akaike Information Criterion (AIC) selected the model K81uf+I+ $\Gamma$ , with base frequencies: A = 0.2972, C = 0.1472, G = 0.1551, T = 0.4005, substitution rate matrix: [A-C] = 1.0000, [A-G] = 201.3902, [A-T] = 35.5380, [C-G] = 35.5380, [C-T] = 201.3902, [G-T] = 1.0000, proportion of invariable sites: (I) = 0.5186, and  $\Gamma$  distribution with the shape parameter 0.7457. The resulting maximum likelihood phylogram (Fig. 2) confirmed placement of *H. dobrogica* within the Cochliopidae, and the close relationships between this species and *H. dalmatica. Heleobops docimus* was placed within the *Heleobia* clade, between *H. dalmatica* and *H. dobrogica*.



**FIGURE 2**. Maximum likelihood phylogram (see text for details). Bootstrap support indicated (10,000 replicates) when > 50%.

## Discussion

Despite all the precautions concerning the molecular clock concept, as well as its scaling (Hillis *et al.* 1996b; Avise 2000; Nei and Kumar 2000; Posada 2003), there are many examples of its usage, also for rissooid snails (Wilke 2003, 2004; Haase *et al.* 2007; Falniowski *et al.* 2007). As a cochliopid, *Heleobia* is phylogenetically not too far from the Hydrobiidae (Wilke *et al.* 2001). The clock was calibrated for two representatives of this family (Wilke 2003). Also the

estimated time (2.172±0.171 Mya) is not far from the 5.33 Mya used as the calibration value. The distances are within the range that is considered not to be affected by saturation (Wilke et al. 2001), and one-point calibration should not give rise to a significant error in this case. However, with onepoint calibration it is not possible to obtain reasonable estimates of confidence intervals (Hillis et al. 1996b). Another problem with calibration was pointed out by Haase et al. (2007). 5.33 Mya is the time of the end of the Messinian Salinity Crisis (Pliocene Flooding). In fact, the isolation of the Atlantic Peringia from the Mediterranean Salenthydrobia must have begun earlier - when the Mediterranean Basin started to separate from the Atlantic, 5.96 Mya (Krijgsman et al. 1999; Falniowski et al. 2007). 5.33 Mya Salenthydrobia became isolated in a freshwater habitat from the other Hydrobia Hartmann, 1821 in the Mediterranean, but its isolation from Peringia Paladilhe, 1874 began earlier. If a time of 5.96 is applied instead of 5.33, the estimates for *Heleobia dobrogica* are higher: 2.429±0.191 Mya (for the Maximum Composite Likelihood distances: 2.669±0.744 Mya – 3.209±0.820 Mya.

Another problem concerns the species we used to estimate the time of divergence. Unfortunately, *Heleobia dalmatica* is the only European cochliopid whose sequence is available. The European *Heleobia* (=*Semisalsa*) is known from the Netherlands, Italy, Croatia, Greece, Romania, Ukraine, Israel, Turkey and Jordan (Kabat and Hershler 1993). Eleven species of this genus have been described, but their status and relationships have not yet been resolved. *Heleobia* (=*Semisalsa*) is probably the only cochliopid representative in the Palearctic, thus its zoogeographic relationships remain enigmatic. According to some authors, *H. dalmatica* is found from Dalmatia to the Black Sea. In any case, both *H. dalmatica* and *H. dobrogica* occur in the Balkans.

Before 3 Mya there was a sharp decrease in temperature and in precipitation. Later, the temperature and humidity became higher, but there were several fluctuations, with alternate periods of cold and warm conditions, and the glacial period in Europe began at about 2.5 Mya (Stanley 1999), predating the Pleistocene. At that time subtropical vegetation definitively disappeared from Europe. The estimated divergence time between these two species, irrespective of the distance used and calibration point assumed, coincides with either the period of climate fluctuation that predated the glaciation period, or the beginning of the glaciation period. It was most probably then that *Heleobia dobrogica* found a safe shelter within a warm cave.

*Heleobia dobrogica* is closely related to *H. dalmatica*, and the K2P distances between these two species suggest that they are congeners (e.g. Wilke *et al.* 2001). Davis *et al.* (1982) synonymized *Semisalsa* Radoman, 1974, with the American *Heleobia* Stimpson, 1865. That was questioned by Bank and Butot (1984). Unfortunately, no COI sequences of any *Heleobia* from South America are available. On the other hand, the American *Heleobops* seems to be very closely related to the European *Heleobia* (=*Semisalsa*).

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